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54) Chemically amplified resist composition.

Positive-tone and negative-tone chemically amplified resist compositions comprising: (a-1) a blocked resin, (a-2) a combination of an alkali-soluble resin and a dissolution controlling agents, or (a-3) a combination of an alkali-soluble resin and a crosslinking agent, (b) a photoacid generator, and (c) specific kinds of solvents. The both positive-tone and negative-tone resist compositions exhibits superior sensitivity, high resolution capability, and excellent storage stability, and can be excellently applied especially to large sized substrates by spincoating for producing excellently shaped patterns by irradiation. The compositions can be used with advantage as a chemically amplified resist for the manufacture of semiconductor devices or integrated circuits (ICs).

The present invention relates to a chemically amplified resist composition, and, more particularly, to a chemically amplified resist composition suitable for super-fin works using radiations, such as ultraviolet rays, deep ultraviolet rays, X-rays, and charged particle rays.

There is an increasing demand for miniaturization of processing sizes in lithography in the fields requiring fin works such as manufacture of semiconductor d vices or integrated circuits (ICs) in order to achieve high integrity in ICs. Technologies capable of processing fine works even of a size of $0.5~\mu m$ or smaller in a stable manner are required in recent years. Because of this, resists used in such a technology must have capability of forming patterns with a size smaller than $0.5~\mu m$ at a high precision. Lithographic techniques using radiations with a shorter wavelength are also being studied.

Included in such radiations are ultraviolet rays (e.g. Hine (365 nm)), deep ultraviolet rays (e.g. KrF excimer laser (248 nm)), X-rays (e.g. synchrotron radiation), and charged particle rays (e.g. electron beam). A number of resists to which these radiations can be applied have been proposed in recent years.

Among such resists, of particular interest are those of which the rate of dissolution in a developer can be changed by an acid produced by irradiation. This kind of resists is generally called "chemically amplified resists" (Chemically Amplified Resists, A. A. Lamola, et al., Lithography Materials & Processing, 53-60, <u>34</u>, No. 8, (1991)).

In the manufacture of ICs using a chemically amplified resist, a resist solution is first prepared. Typically, this resist solution comprises (a) a resin protected by a group decomposable by an acid (hereinafter referred to as "acid-decomposable protective group"), which is converted into an alkali-soluble resin when the acid-decomposable protective group is decomposed (such a protected resin is more specifically defined hereinafter and referred to as "blocked resin"), (b) a compound which generates an acid by irradiation (hereinafter referred to as "photoacid generator"), and (c) a solvent which dissolves the blocked resin and the photoacid generator. This resist solution is then applied to a substrate by using a spin coater or a roll coater to form a resist coating. Adequate coatability and storage stability are thus essential characteristics for the resist solutions in order to enable sophisticated fine works to be constantly implemented. Irradiation to the resist coating produces patterns suitable for fine works. The shape of the patterns greatly affects the precision of the fine works, and a rectangular pattern is considered to be a preferred pattern.

Ethylene glycol monoethylether acetate and the like are known as solvents used for the preparation of a conventional lithographic resist solution comprising a novolak resin and a naphthoquinone diazide compound as a photosensitizer. The use of ethylene glycol monoethylether acetate as a solvent for preparing a solution of a chemically amplified resist poses a problem of storage instability. Depending on the period of time for which the resist solution prepared from this solvent has been stored, there may occur unevenness in the resist coating sensitivity, resulting in irregular patterns.

Further, along with the requirements for high integrity of ICs in recent years, substrates (silicone wafers) with a larger diameter, e.g., 4 inch, 6 inch, or even 8 inch, are employed in order to increase the yield and to promote the efficiency in the manufacture of ICs. A solution of chemically amplified resist prepared using ethylene glycol monoethylether acetate as a solvent exhibits only inadequate spin-coatability on such large diameter substrates.

An object of the present invention is therefore to provide a novel composition for chemically amplified resist which exhibits superior sensitivity and resolution capability and can be excellently applied especially to large sized substrates by spincoating.

Another object of the present invention is to provide a composition for chemically amplified resist which exhibits excellent storage stability and forms resist coating for producing excellently shaped patterns, thereby ensuring fine works to be constantly implemented.

Still another object of the present invention is to provide a composition for chemically amplified resist having superior storage stability, which can exhibit high sensitivity and produce excellently shaped patterns with good reproducibility even after storage for a long period of time.

The present inventors have undertaken extensive studies, and found that the use of specific kinds of solvents in combination with (a-1) a blocked resin, (a-2) a combination of an alkali-soluble resin and a dissolution controlling agents (hereinafter defined), or (a-3) a combination of an alkali-soluble resin and a crosslinking agent, and (b) a photoacid generator can achieve the above objects.

The present inventors have further found that the following solvents can be excellently used as said specific kinds of solvents for chemically amplified resist compositions of negative-tone type and of positive-tone type.

- (1) Solvents used for negative-ton type amplified resist compositi ns:
 - (i) an ester of a saturated carboxylic acid having 1-4 carbon atoms with no substituent and an alcoh I having 1-6 carbon atoms (hereinafter referred to as "ester solv nt"),
 - (ii) a mixture of ester solvent (i) and an alkyl lactate, or

- (iii) a mixture of an alkyl lactate, a propyl ne glycol alkyl ether and/ r a propyl ne glycol alkyl ther acetat.
- (2) Solvents used for positive-tone type amplified resist compositi ns:
 - (i) the ester solvent,
 - (ii) a mixture of ster solv nt (i) and an alkyl lactate,
 - (iii) a mixture of an alkyl lactate, a propylene glycol alkyl ether and/or a propylene glycol alkyl ether acetate. or
 - (iv) a propylene glycol alkyl ether acetate.

Accordingly, one of the specific objects of the present invention is to provide a positive-tone type chemically amplified resist composition (hereinafter referred to as "positive-tone resist composition") comprising,

- (A) a resin of which the acidic functional groups are protected by an acid decomposable group and which is converted into an alkali-soluble resin when the acid-decomposable protective group is decomposed (a blocked resin),
- (B) a compound which generates an acid by radiation (photoacid generator), and
- (C) a solvent selected from the group consisting of,
 - (i) an ester solvent,

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- (ii) a mixture of ester solvent (i) and an alkyl lactate,
- (iii) a mixture of an alkyl lactate, a propylene glycol alkyl ether and/or a propylene glycol alkyl ether acetate, and
- (iv) a propylene glycol alkyl ether acetate.

Another specific objects of the present invention is to provide a positive-tone resist composition comprising,

- (A') an alkali-soluble resin,
- (D) a compound which inhibits dissolution of the alkali-soluble resin in an alkali and is decomposable in the presence of an acid, and, when decomposed in the presence of an acid, exhibits an action of reducing or losing the alkali insolubility of a resin coating, or accelerating dissolution of the alkali-soluble resin in alkali (hereinafter referred to as "dissolution controlling agent"),
 - (B) a photoacid generator, and
 - (C) a solvent selected from the group consisting of said solvents (i)-(iv).

Still another specific object of the present invention is to provide a negative-tone type chemically amplified resist composition (hereinafter referred to as "negative-tone resist composition") comprising,

- (A') an alkali-soluble resin,
- (E) a compound which can crosslink the alkalisoluble resin in the presence of an acid (hereinafter referred to as "crosslinking agent"),
 - (B) a photoacid generator, and
 - (C) a solvent selected from the group consisting of,
 - (i) an ester solvent,
 - (ii) a mixture of ester solvent (i) and an alkyl lactate, and
 - (iii) a mixture of an alkyl lactate, a propylene glycol alkyl ether and/or a propylene glycol alkyl ether acetate.

These and other objects, features, and advantages of the present invention will become more apparent from the following description of the preferred embodiments.

Each component of the chemically amplified compositions of the present invention are hereinafter discussed in detail.

45 Alkali-soluble resins (Component (A'))

There are no specific limitations to the alkalisoluble resins used in the positive-tone resist composition or the negative-tone resist composition, so long as such resins have a functional group exhibiting affinity to an alkali developer, such as acidic functional groups (e.g. phenolic hydroxy group or carboxy group), and are soluble in an alkali developer.

Typical examples of such alkali-soluble resins are vinyl polymers containing a repeating unit formed by the cleavage of a polymerizable double bond of a monomer having at least one acidic functional group, such as hydroxystyrene, hydroxy-α-methylstyrene, vinylbenzoic acid, carboxymethylstyrene, carboxymethoxystyrene, (meth)acrylic acid, crotonic acid, maleic acid, itaconic acid, citraconic acid, mesaconic acid, or cinnamic acid, and condensed polymers containing a condensed repeating unit having an acidic functional group, typified by novolak resins.

In the cas where said vinyl polymer is used as the alkali-soluble resin, although such resin may consist only of said repeating unit formed by thi cleavag of a polymerizable double bond of a monomer having at

least on acidic functional group, other repeating units may b contain d as need d, inasmuch as the resin is soluble in an alkali developer.

Examples of such ther repeating units include those formed by thicl avag of thipolymerizable double bonds derived from monomers such as styrene, α-methylstyrene, vinyltoluene, malic anhydride, (meth)acryl nitrile, crotonitrile, maleinitrili, fumaronitrile, mesaconitrile, citraconitrile, itaconitrili, (meth)acryamide, crotonamide, maleinamide, fumaroamide, mesaconamide, citraconamide, itaconamide, vinylaniline, vinylpyridien, vinyl-ε-caprolactum, vinylpyrrolidon, and vinylimidazole.

The polymerization or copolymerization reactions for manufacturing said alkali-soluble vinyl polymers can be carried out by bulk polymerization, solution polymerization, precipitation polymerization, emulsion polymerization, suspension polymerization, or bulk-suspension polymerization, in which polymerization initiators (e.g., radical polymerization initiators, anionic polymerization initiators, conjugated anionic polymerization initiators, cationic polymerization initiators) and polymerization catalysts are suitably selected depending on the kinds of monomers and reaction media.

In the case where said condensed polymer is used as the alkali-soluble resin, although it is possible that such a resin consists only of said repeating unit such as novolak resin moieties, other repeating units may be additionally contained, inasmuch as the resulting resin is soluble in an alkali developer. Such condensed polymers can be manufactured by the polycondensation or copolycondensation of at least one phenol compound and at least one aldehyde compound, optionally, together with other polycondensable compounds, in an aqueous medium or in a mixture of water and a solvent having an affinity with water in the presence of an acid catalyst.

Here, given as examples of phenol compounds are o-cresole, m-cresole, p-cresole, 2,3-xylenol, 2,4-xylenol, 2,5-xylenol, 3,5-xylenol, 3,5-xylenol, 2,3,5-trimethylphenol, and 3,4,5-trimethylphenol; and as aldehyde compounds, formaldehyde, trioxane, paraformaldehyde, benzaldehyde, acetaldehyde, propylaldehyde, and phenylacetaldehyde.

The amount of the repeating units having an acidic functional group in the alkali-soluble resin cannot be generically determined, because such an amount varies depending on the other repeating units which are optionally incorporated. Typically, this amount is 15-100 mol%, and preferably 20-100 mol%.

The polystyrene-reduced weight average molecular weight of the alkali-soluble resin measured by gel permeation chromatography (hereinafter referred to as "Mw") is preferably 1,000-150,000, and more preferably 3,000-100,000.

The alkali-soluble resins may be hydrogenated in the case where they contain repeating units having unsaturated carbon-carbon bonds. The degree of hydrogenation to saturate 70% or less, preferably 50% or less, and more preferably 40% or less, of unsaturated carbon-carbon bonds contained in such recurring units is sufficient. If the hydrogenation degree exceeds 70%, the developing characteristics may be impaired when the alkali-soluble resins are developed with an alkaline developer.

The alkali-soluble resins can be used either alone or in combination of two or more of them in the positive-tone resist composition or the negative-tone resist composition.

Blocked resin (Component (A))

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The blocked resin in the present invention is the resin having acidic functional groups (e.g. phenolic hydroxy group or carboxy group) in the alkali-soluble resins blocked by one or more groups which are decomposable in the presence of an acid (acid-decomposable protective group). The blocked resins are thus insoluble or scarcely soluble in alkali, but become alkali-soluble when the acid-decomposable protective group is decomposed by an acid generated from a photoacid generator. The term "insoluble or scarcely soluble in alkali" is defined as the characteristics of a resin, in which at least 50% (in thickness) of the coating prepared from the resin remains undissolved when developed under the same alkaline developing conditions as the conditions under which resist patterns are formed on a resist coating made from the positive-tone or negative-tone resist composition.

The acid-decomposable protective group is a group which protects or blocks the acidic functional groups, such as phenolic hydroxy group or carboxy group, and which is decomposed in the presence of an acid. There are no specific restrictions to the acid-decomposable protective groups so long as these properties are satisfied, and such groups include, for example, substituted methyl groups, 1-substituted ethyl groups, 1-branched alkyl groups, silyl groups, germyl groups, alkoxycarbonyl groups, acyl groups, and cyclic acid-decomposable protective groups.

Giv n as examples of said substituted methyl groups are m thoxymethyl group, m thylthiomethyl group, ethoxymethyl group, ethoxym thyl group, benzylthiomethyl group, benzylthiomethyl group, benzylthiomethyl group, benzylthiophenacyl group, methoxyph nacyl group, (methylthio)phenacyl group,

cycl propylmethyl group, benzyl group, diphenylmethyl group, triphenylmethyl group, bromobenzyl group, nitrobenzyl group, methoxybenzyl group, methylthiobenzyl group, thoxybenzyl group, thylthiobenzyl group, and piperonyl group.

Given as examples of said 1-substituted ethyl groups are 1-meth xyethyl group, 1-methylthioethyl group, 1,1-dimethoxy thyl group, 1-ethoxy thyl group, 1-ethylthio thyl group, 1,1-di thoxyethyl group, 1-phenoxyethyl group, 1-phenylthioethyl group, 1,1-diphenoxyethyl group, 1-benzylthioethyl group, 1-benzylthioethyl group, 1-cyclopropylethyl group, 1-phenylethyl group, 1,1-diphenylethyl group, and α-methylphenacyl group.

Included in said 1-branched alkyl groups are, for example, isopropyl group, sec-butyl group, t-butyl group, 1,1-dimethylpropyl group, 1-methylbutyl group, and 1,1-dimethylputyl group.

Examples of said silyl groups are trimethyl silyl group, ethyldimethyl silyl group, diethylmethyl silyl group, triethyl silyl group, dimethyl-iso-propyl silyl group, triethyl silyl group, dimethyl-iso-propyl silyl group, tri-t-butyl silyl group, dimethyl silyl group, dimethyl silyl group, methyl-diphenyl silyl group, and triphenyl silyl group.

Examples of said germyl groups are trimethyl germyl group, ethyldimethyl germyl group, diethylmethyl germyl group, triethyl germyl group, dimethyl-iso-propyl germyl group, methyl-di-iso-propyl germyl group, tri-iso-propyl germyl group, t-butyldimethyl germyl group, di-t-butylmethyl germyl group, tri-t-butyl germyl group, dimethylphenyl germyl group, methyldiphenyl germyl group, and triphenyl germyl group.

Given as examples of said alkoxycarbonyl groups are methoxycarbonyl group, ethoxycarbonyl group, isopropoxycarbonyl group, t-buthoxycarbonyl group, and t-pentyloxycarbonyl group.

Further, given as examples of acyl groups are acetyl group, propionyl group, butyryl group, heptanoyl group, hexanoyl group, valeryl group, pivaloyl group, isovaleryl group, lauryloyl group, myristoyl group, palmytoyl group, stearoyl group, oxalyl group, malonyl group, scucinyl group, glutaryl group, adipoyl group, piperoyl group, suberoyl group, azelaoyl group, sebacoyl group, acryloyl group, propioyl group, methacryloyl group, crotonoyl group, oleoyl group, maleoyl group, fumaroyl group, mesaconoyl group, campholoyl group, benzoyl group, phthaloyl group, iso-phthaloyl group, terephthaloyl group, naphthoyl group, toluoyl group, hydroatropoyl group, atropoyl group, cinnamoyl group, furoyl group, thenoyl group, nicotinoyl group, isonicotinoyl group, toluvenesulfonyl group, and mesyl group.

Given as examples of said cyclic acid-decomposable protective groups are cyclopropyl group, cyclopentyl group, cyclohexyl group, cyclohexyl group, tetrahydropyranyl group, 4-methoxycyclohexyl group, tetrahydropyranyl group, tetrahydrothiopyranyl group, tetrahydrothiofuranyl group, 3-bromote-trahydropyranyl group, 4-methoxytetrahydropyranyl group, 4-methoxytetrahydropyranyl group, 2-1,3-dithiolanyl group, benzo-2-1,3-dioxolanyl group, and benzo-2-1,3-dithiolanyl group.

Among these acid-decomposable protective groups, especially preferred are t-butyl group, benzyl group, t-butoxy-carbonyl group, tetrahydropyranyl group, tetrahydrofuranyl group, tetrahydrothiopyranyl group, and tetrahydrothiofuranyl group.

The amount of the acid-decomposable protective groups to be introduced to the blocked resins (the percentage of the number of the acid-decomposable protective groups in the total number of the acidic functional groups and the acid-decomposable protective groups) is preferably 15-100%, more preferably 20-100%, and particularly preferably 20-80%.

The Mw of the blocked resin is preferably 1,000-150,000, and more preferably 3,000-100,000.

The blocked resin can be prepared, for example, by introducing at least one acid-decomposable protective group to alkali-soluble resin; by the polymerization or copolymerization of monomer containing at least one acid-decomposable protective group; or by the polycondensation or copolycondensation of polycondensable component containing at least one acid-decomposable protective group.

These blocked resins used in the positive-tone resist composition not only are made alkali-soluble themselves by an acid generated from a photoacid generator when the composition is irradiated, but also possess a characteristic of controlling the solubility of alkali-soluble resins in an in alkali solution. That is to say, these resins can suppress the solubility of alkali-soluble resins in alkali, rendering the resin coating alkali-insoluble and, when decomposed in the presence of an acid, can exhibit an action of reducing or losing the alkali-insolubility of the resin coating, or an action of accelerating dissolution of the alkalisoluble resin in an alkali. The blocked resins can thus be used also as a dissolution controlling agent (Component (D)) for the alkali-soluble resin (A') in one embodiment of the positive-tone resist composition of the present invention.

These blocked resins may be used either alone or in combination of two or more of them.

Dissolution controlling ag nt (Component (D))

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The dissolution controlling agent which is used in combination with the alkali-soluble resin (A') in a embodiment of the positive-tone resist composition of the present invention is a compound which inhibits dissolution of the alkali-soluble resin in an alkali and is decomposable in the presence of an acid, and, when decomposed in the presence of an acid, exhibits an action of reducing or losing the alkali insolubility of a resin coating, or accelerating dissolution of the alkalisoluble resin in alkali.

Compounds having an acid-decomposable protective group substituted for acidic functional groups such as phonolic hydroxy group or carboxy group can be given as examples of the dissolution controlling agent. The above-mentioned blocked resins are also included in the dissolution controlling agents.

The dissolution controlling agent may be either a low molecular weight compound or a high molecular weight compound. Preferable dissolution controlling agents are compounds obtained by introducing said acid-decomposable protective groups to a poly-phenolic compound (e.g. bisphenol A, bisphenol F, or bisphenol S) or to a carboxylic acid compound (e.g. hydroxyphenylacetic acid).

Compounds of the following formulas (1) and (2) are examples of particularly preferable dissolution controlling agents.

$$t-C_4H_9-OCO - CH_3 - OCO - t-C_4H_9$$

$$CH_3 - OCO - t-C_4H_9$$

$$CH_3 - OCO - t-C_4H_9$$

$$CH_3 - OCO - t-C_4H_9$$

The above-described blocked resins (Component (A)) can be used as the dissolution controlling agent of a high molecular weight compound.

These low molecular weight compounds and high molecular weight compounds can be used as the dissolution controlling agent in the positive-tone resist composition either alone or in combination of two or more of them. The combination of the low molecular weight compound and the high molecular weight compound is also possible.

Crosslinking agents (Component (E))

The crosslinking agent used in the negative-tone resist composition is a compound which can crosslink alkali-soluble resins in the presence of an acid, for example, the acid generated by radiation. As an example of such a crosslinking agent, a compound having at least one substituent which exhibits crosslinking reactivity with the alkali-soluble resin can be given (such a group is hereinafter referred to as "crosslinkable substituent").

The following groups (3)-(7) are given as specific examples of the crosslinkable substituents.

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$$--X - CH_2 - C$$

wherein 1 is 1 or 2; X represents, when 1 is 1, a single bond -O-, -S-, -CO-O-, or -NH-, and when 1 is 2, a trivalent N; Y is -O- or -S-; m is 0 or an integer of 1 to 3; and n is an integer of 1 to 3, provided that n + m is 4 or smaller.

$$--\left(C(R^{1})(R^{2})\right)_{k}Z--R^{3}$$
(4)

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wherein k is 0 or an integer of 1 or more; Z represents -O-, -CO-O-, or -CO-; R¹ and R², which may be the same or different, represent a hydrogen atom or an alkyl group having 1-4 carbon atoms; and R³ represents an alkyl group having 1-5 carbon atoms, an aryl group having 6-12 carbon atoms, or an aralkyl group having 7-14 carbon atoms.

$$-C(R^4)=C(R^5)(R^6)$$
 (5)

5 wherein R⁴, R⁵ and R⁶, which may be the same or different, represent a hydrogen atom or an alkyl group having 1-4 carbon atoms.

35 wherein k is 0 or an integer of 1 or more; R¹ and R², which may be the same or different, represent a hydrogen atom or an alkyl group having 1-4 carbon atoms; and R³ and R³ each individually represent an alkylol or an alkoxyalkyl group having 1-5 carbon atoms.

$$\frac{-\left(C(R^1)(R^2)\right)}{k}N R^9 \tag{7}$$

wherein k is 0 or an integer of 1 or more; R¹ and R², which may be the same or different, represent a hydrogen atom or an alkyl group having 1-4 carbon atoms; and R³ is a divalent organic group which can form 3- to 8-membered ring containing a hetero atom selected from oxygen, sulfur and nitrogen.

Given as specific examples of these crosslinkable substituents are glycidyl ether group, glycidyl ester group, glycidyl amino group, methoxymethyl group, ethoxymethyl group, benzyloxymethyl group, diethylaminomethyl group, dimethylaminomethyl group, diethylolaminomethyl group, morpholinomethyl group, acetoxymethyl group, benzoyloxymethyl group, formyl group, acetyl group, vinyl group, and isopropenyl group.

Specific examples of the compounds having these crosslinkable substituents are epoxy compounds of bisphenole A, epoxy compounds of bisphenole F, epoxy compounds of bisphenole S, novolak resin-type epoxy compounds, resoleresin-type poxy compounds, poly(hydroxystyren)-type epoxy compounds, methylole group-containing metamine compounds, methylole group-containing benzoguanamine compounds, methylole group-containing phenolecompounds, alkoxyalkylegroup-containing phenolecompounds, alkoxyalkylegroup-containing urea compounds.

containing melamine compounds, alkoxyalkyl group-containing benzoguanamine compounds, alkoxyalkyl group-containing urea compounds, alkoxyalkyl group-containing phenol compounds, carboxymethyl group-containing melamine compounds, carboxymethyl group-containing benzoguanamine compounds, carboxymethyl group-containing phen I compounds.

Among these, methylol group-containing phenol compounds, methoxym thyl group-containing melamin compounds, methoxymethyl group-containing phenol compounds, and acetoxymethyl group-containing phenol compounds are particularly preferred, with an ideal compound being methoxymethyl group-containing melamine compounds. CYMEL 300, CYMEL 301, CYMEL 303, and CYMEL 305 (trademarks, manufactured Mitsui Cyanamid Co.) are given as examples of commercially available methoxymethyl group-containing melamine compounds. Chemical structure of the typical compound is shown by the following formula (8).

Further, the alkali-soluble resins afforded with crosslinking characteristics by introducing one of said crosslinkable substituents to acidic functional group thereof are suitable crosslinking agents. The proportion of crosslinkable substituents introduced to the alkalisoluble resin is normally 5-60 mol%, preferably 10-50 mol%, and more preferably 15-40 mol%. If this proportion is less than 5 mol%, it is difficult to effect a satisfactory crosslinking reaction, resulting in reduction in the rate of residual coatings and inducing winding or swelling of patterns; if more than 60 mol%, solubility of the alkali-soluble resin is decreased, resulting in impaired developing performances.

These crosslinking agents can be used either alone or in combination of two or more of them in the negative-tone resist composition.

Photoacid generators (Component (B))

Included in the photoacid generators, which are compounds generating an acid by irradiation, are onium salts, halogen-containing compounds, sulfones, and sulfonates. Specific examples of these compounds are as follows.

(a) Onium salts

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lodonium compounds, sulfonium compounds, phosphonium compounds, diazonium compounds, pyridium compounds, and the like are given. Of these preferred compounds are diphenyliodonium trifluoromethanesulfonate, diphenyliodonium pyrenesulfonate, diphenyliodonium dodecylbenzenesulfonate, triphenylsulfonium trifluoromethanesulfonate, triphenylsulfonium hexafluoroantimonate, triphenylsulfonium naphthalenesulfonate, (hydroxyphenyl)benzylmethylsulfonium toluenesulfonate, and the like.

(b) Halogen-containing compounds

Haloalkyl group-containing heterocyclic compounds, haloalkyl group-containing hydrocarbon compounds, and the like are given as typical examples. Preferred specific compounds include derivatives of (trichlorormethyl)-s-triazine, such as phenyl-bis(trichlorormethyl)-s-triazine, methoxyphenyl-bis(trichlorormethyl)-s-triazine, and naphtyl-bis(trichlorormethyl)-s-triazine; and 1,1-bis(4-chlorophenyl)-2,2,2-tri-chloroethane.

(c) Sulfone compounds

 β -ketosulfone, β -sulfonylsulfone, and α -diazo compounds of these are given as typical examples. Preferred compounds are phenacylphenyl sulfone, methylphenacyl sulfone, bis(phenylsulfonyl)methane, and bis(phenylsulfonyl)diazomethane.

(d) Sulfonate compounds

Alkylsulfonates, haloalkylsulfonates, arylsulfonates, and iminosulfonates are given as examples. Preferred compounds include benzointosylate, tris-triflu romethanesulf nate of pyrogallol, nitrobenzyl-9,10-dieth xyanthracene-2-sulfonate, and the lik.

These photoacid generators may be used either alon or in combination of two or more of th m.

Solvents Compon nt (C))

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The solvents used in the chemically amplified resist compositi ns of the present inv ntion are as follows.

- (1) For the negative-tone resist composition a solvent selected from the following groups are used.
 - (i) An ester of a saturated carboxylic acid having 1-4 carbon atoms with no substituent and an alcohol having 1-6 carbon atoms (ester solvent).
 - (ii) A mixture of ester solvent (i) and an alkyl lactate.
 - (iii) A mixture of an alkyl lactate, a propylene glycol alkyl ether and/or a propylene glycol alkyl ether acetate
- (2) For positive-tone resist compositions a solvent selected from the following groups are used.
 - (i) Said ester solvent.
 - (ii) A mixture of ester solvent (i) and an alkyl lactate.
 - (iii) A mixture of an alkyl lactate, a propylene glycol alkyl ether and/or a propylene glycol alkyl ether acetate.
 - (iv) A propylene glycol alkyl ether acetate.

These solvents afford excellent storage stability to the chemically amplified resist compositions. The chemically amplified resist compositions using these solvents can exhibit superior sensitivity and produce fine resist patterns even after having been stored for a long period of time.

Solvent (i)

Solvent (i) is an ester solvent prepared by condensation of a saturated carboxylic acid having 1-4 carbon atoms with no substituent and an alcohol having 1-6 carbon atoms.

Examples of the saturated carboxylic acid having 1-4 carbon atoms with no substituent include formic acid, acetic acid, propionic acid, and butyric acid. Given as examples of the alcohol having 1-6 carbon atoms are methanol, ethanol, n-propanol, iso-propanol, n-butanol, sec-butanol, tert-butanol, amyl alcohol, iso-amyl alcohol, hexanol, and cyclohexanol.

Included in the ester solvents prepared from these carboxylic acids and alcohols are methyl formate, ethyl formate, amyl formate, methyl acetate, ethyl acetate, propyl acetate, iso-propyl acetate, n-butyl acetate, iso-butyl acetate, amyl acetate, iso-amyl acetate, hexyl acetate, methyl propionate, ethyl propionate, n-butyl propionate, methyl butylate, ethyl butylate, propyl butylate, iso-propyl butylate, n-butyl butylate, iso-butyl butylate, and the like.

Of these preferred ester solvents are amyl formate, methyl acetate, ethyl acetate, n-butyl acetate, iso-butyl acetate, iso-amyl acetate, ethyl n-butylate, iso-propyl butylate, and n-butyl butylate.

Solvent (ii)

As preferable alkyl lactates which can be used mixed with said ester solvent (i), C_1 - C_8 alkyl lactates, such as methyl lactate, ethyl lactate, and butyl lactate, can be given. Especially preferred is ethyl lactate.

The ester solvent and alkyl lactate are preferably incorporated at a ratio of 30-90:70-10 parts by weight in 100 parts by weight of solvent (ii).

5 Solvent (iii)

Solvent (iii) comprises said alkyl lactate and a propylene glycol alkyl ether and/or a propylene glycol alkyl ether acetate. Included in propylene glycol alkyl ethers which can be used here are propylene glycol monoalkyl ethers and propylene glycol dialkyl ethers.

There are two isomers for propylene glycol; 1,2-dihydroxy compound and 1,3-dihydroxy compound. Further, are there are two isomers for monoalkyl ether of 1,2-dihydroxy compound. Thus, there are three isomers for each propylene glycol monoalkyl ether. Each propylene glycol dialkyl ether also has three isomers, with the exception where the two alkyl groups are the same, in which case the number of isomers are two. All these isomers can b used as components of solv nt (iii).

Lower-alkyl ethers having 1-6 carbon atoms are preferably used as the alkyl ether group for forming the propylene glycol alkyl ether. Preferred alkyl ethers are methyl ther, ethyl ther, propyl ether, butyl ther, and the like, with methyl ether and ethyl ether being particularly preferred.

Specific examples of propyl ne glycol alkyl ethers are propylene glycol monomethyl ther, propyl ne gly-

col monoethyl ether, propylene glycol monopropyl ether, propylene glycol monobutyl ether, propylene glycol dimethyl ther, propylene glycol dibutyl ether, propylene glycol dibutyl ether. Particularly preferred propylene glycol alkyl ethers are propyl ne glycol monomethyl ther, propylene glycol dimethyl ether, and propyl ne glycol dibutyl ether.

In the present invintion propyl ne glycol monoalkyl ethers and propyl ne glycol dialkyl thers, including isomers and those having different alkyl ether groups, may be used either alone or in combination of two or more of them. Also, a mixture of a mono-ether compound and di-ether compound can be preferably used.

Now, illustrating propylene glycol alkyl ether acetates used as a component for solvent (iii), in the same manner as in the case of propylene glycol alkyl ethers, there are two isomers for propylene glycol, which are 1,2-dihydroxy compound and 1,3-dihydroxy compound. Further, there are two isomers for monoalkyl ether acetate of 1,2-dihydroxy compound. Thus, there are three isomers for each propylene glycol monoalkyl ether acetate.

The same lower-alkyl ethers having 1-6 carbon atoms as given for the propylene glycol alkyl ethers can be preferably used as the alkyl ether group for forming the propylene glycol alkyl ether acetates. Preferred alkyl ethers are methyl ether, ethyl ether, propyl ether, butyl ether, and the like, with methyl ether and ethyl ether being particularly preferred.

Preferred examples of propylene glycol alkyl ether acetates are lower-alkyl (C₁-C₆) ether acetates of propylene glycol, such as propylene glycol monomethyl ether acetate, propylene glycol monomethyl ether acetate, and propylene glycol monomethyl ether acetates, with propylene glycol monomethyl ether acetate being most preferred compound.

These propylene glycol monoalkyl ether acetates, irrespective of the kinds of the alkyl ether portion or the kinds of the isomers, may be used either alone or in combination of two or more of them.

For 100 parts by weight of solvent (iii), alkyl lactate is preferably contained 30-95 parts by weight, with the remaining portion being propylene glycol alkyl ether and/or propylene glycol alkyl ether acetate.

Solvent (iv)

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Beside the above-described solvents (i) to (iii), propylene glycol alkyl ether acetate can be used independently for positive-tone resist compositions. All illustrations concerning the propylene glycol alkyl ether acetates in solvent (iii) above are applicable relating to the propylene glycol alkyl ether acetates of solvent (iv).

Beside the above-described solvents (i) to (iv) which are essential in the chemically amplified resist composition of the present invention, other solvents can be used to the extent of 70 parts by weight, preferably 50 parts by weight, and more preferably 30 parts by weight, in 100 parts by weight of the solvents.

Included in such other solvents which can be used are ethylene glycol monoalkyl ethers, such as ethylene glycol monomethyl ether, ethylene glycol monopropyl ether, and ethylene glycol monopropyl ether, and ethylene glycol monomethyl ether acetates, such as ethylene glycol monomethyl ether acetate and ethylene glycol monoethyl ether acetate; diethylene glycol dialkyl ethers, such as diethylene glycol dimethyl ether, diethylene glycol dipropyl ether, and diethylene glycol dibutyl ether; aromatic hydrocarbons, such as toluene and xylene; ketones, such as methyl ethyl ketone, 2-heptanone, 3-heptanone, 4-heptanone, and cyclohexanone; esters, such as ethyl 2-hydroxy-2-methylpropionate, ethyl ethoxyacetate, ethyl hydroxyacetate, methyl 2-hydroxy-3-methylbutyrate, methyl 3-methoxypropionate, ethyl 3-ethoxypropionate, and ethyl 3-methoxypropionate, 3-methyl-3-methoxybutylacetate, 3-methyl-3-methoxybutylacetate, ethyl acetoacetate, ethyl acetoacetate, methyl pyruvate, and ethyl pyruvate; amides, such as N-methylpyrrolidone, N,N-dimethylformamide, N-methylformamide, and N,N-dimethylacetamide.

These other solvents may be used either alone or in combination of two or more of them.

Proportions of components in the compositions

The proportions of the above-described components to be incorporated in the chemically amplified resist compositions of the present invention are as follows.

The proportion of photoacid generator incorporated in the positive-tone resist composition or the negative-tone resist composition is preferably 0.05-20 parts by weight, more preferably 0.1-15 parts by weight, and particularly preferably 0.5-10 parts by weight, for 100 part by weight of the alkali-soluble resin and/or the blocked resin. If the proportion of the photoacid generators is less than 0.05 parts by weight, it may be difficult for the acid-catalyzed chemical reaction induced by the acid produced by irradiation to effectively take place. If it is greater than 20 parts by weight, unevenues in the coating may occur when the positive-tone resist composition or the negative-tone resist composition is applied, and there may be the occasient where scums are pro-

duced during development.

When the combination of the dissolution controlling agent and the alkali-soluble resins is used for the positive-tone resist composition, the dissolution controlling agents are incorporated preferably in an amount of 5-150 parts by weight, more preferably 5-100 parts by weight, and particularly preferably 5-50 parts by weight for 100 part of the alkali-soluble resins.

If the amount of the dissolution controlling agents is smaller than 5 parts by weight, desired effects are exhibited only with difficulty; if it is greater than 150 parts by weight, there is a tendency that coatability of the positive-tone resist composition is decreased, producing coating with poor strength.

The crosslinking agents in the negative-tone resist composition are incorporated preferably in an amount of 5-95 parts by weight, more preferably 15-85 parts by weight, and particularly preferably 20-75 parts by weight for 100 parts by weight of the alkali-soluble resins. If this proportion of the crosslinking agents is less than 5 parts by weight, it is difficult to effect a satisfactory crosslinking reaction, resulting in reduction in the rate of residual coating and inducing winding or swelling of patterns; if more than 95 parts by weight, production of scums may increase, resulting in impaired development.

The amounts of solvents used in the positive-tone resist composition or the negative-tone resist composition is normally 20-3,000 parts by weight, preferably 50-3,000 parts by weight, and more preferably 100-2,000 parts by weight, for 100 parts by weight of the total amount of alkali-soluble resins and/or the blocked resins.

Other additives

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Various additives, such as surface active agents and sensitizers, can be incorporated in the chemically amplified resist composition of the present invention as needed.

Surface active agents improve coatability and promote developing performances of the chemically amplified resist. Included in such surface active agents are polyoxyethylene lauryl ether, polyoxyethylene stearyl ether, polyoxyethylene oleyl ether, polyoxyethylene octylphenyl ether, polyoxyethylene nonylphenyl ether, polyoxyethylene glycol dilaurate, and polyoxyethylene glycol distearate, as well as commercially available surfactants, such as KP341 (trademark, manufactured by Sin-etsu Chemical), Polyflow No. 75, No. 95 (trademarks, manufactured by Kyoei Oil and Fat Chemical), Efftop EF301, EF303, EF352 (trademarks, manufactured by Tokem Products), Megafack F171, F172, F173 (trademarks, manufactured by Dainippon Ink and Chemicals), Florade FC430, FC431 (trademarks, manufactured by Surnitomo 3M), Asahi Guard AG710, Surflon S-382, SC-101, SC102, SC-103, SC-104, SC-105, SC-106 (trademarks, manufactured by Asahi Glass), and the like.

The amount of the surface active agents to be incorporated is normally less than 2 parts by weight for 100 parts by weight of the chemically amplified resist composition.

Sensitizers absorb irradiated energy and transfer the absorbed energy to the photoacid generators, thereby accelerating production of an acid. Thus, the sensitizers have an effect of increasing the sensitivity of resist coating made from the chemically amplified resist composition. Preferable examples of such sensitizers are acetone, benzene, acetophenone, benzophenoene, naphthalene, biacetyl, eosine, Rosebengal, pyrenes, anthracenes, phenothiazines, and the like.

The amount of the sensitizers to be incorporated is normally less than 50 parts by weight, preferably 30 parts by weight, for 100 parts by weight of the chemically amplified resist composition.

Further, dyes or pigments may be incorporated for alleviating effects of halations during irradiation, and adhesion promoters may be added to improve adhesion properties of the resist coating to substrates.

Furthermore, other additives, such as halation inhibitors (e.g., azo compounds, amine compounds), preservatives, defoaming agents, shape improvers, and the like may be added to the chemically amplified resist composition of the present invention.

Preparation of the compositions

The chemically amplified resist composition of the present invention can be prepared by dissolving various solid components in the solvent to obtain a solution containing 5-50% by weight of solid components, filtering the solution through a filter with approximately $0.2 \mu m$ pore size.

Preparation of resist pattern

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In rider to prepare a resist pattern using the chemically amplified resist composition of the present invention, the composition is applied to a substrate such as a lilicon wafer, aluminum-coated silicon wafer, to form a resist coating, and this resist coating is irradiated to form a desired pattern. Radiation used here is suitably

selected from ultraviolet rays (e.g. i-lin), deep ultraviolet rays (e.g. excimer laser), X-rays (.g. synchrotron radiation), and charged particle rays (e.g. electron beam). Irradiati n conditions such as the exposure dose of irradiation are appropriately determined d pending n the kind of th chemically amplified resist composition.

When a resist pattern is prepared using the chemically amplified resist composition of the present invinction, a protective coating may be provided over the resist coating in order to prevent the effects of impurities such as basic impurities contained in the working atmosphere.

Further, it is desirable for the resist coating to be baked after the irradiation in order to improve its apparent sensitivity (post-exposure bake). The heating conditions of baking is typically 30-200°C, and preferably 50-150°C, although the temperature may be varied depending on the proportion of components in the composition and kinds of additives.

Then, the resist coating is developed with an alkaline developer for forming a prescribed pattern. An alkaline solution in which an alkaline compound is dissolved to a concentration of 1-10% by weight, preferably 2-5% by weight, is used as the alkaline developer. Given as examples of alkaline compounds which can be used here are sodium hydroxide, potassium hydroxide, sodium carbonate, sodium silicate, sodium methasilicate, ammonia water, ethylamine, n-propylamine, diethylamine, di-n-propylamine, triethylamine, methyldiethylamine, dimethylethanolamine, triethanolamine, tetramethylammonium hydroxide, tetraethylammonium hydroxide, choline, pyrole, piperidine, 1,8-diazabicyclo-[5,4,0]-7-undecene, and 1,5-diazabicyclo-[4,3,0]-5-non-ene.

Further, suitable amounts of water-soluble organic solvents, such as methanol and ethanol, and surface active agents may be added to the developer.

The developed products are normally washed with water after development using these alkaline developers.

Japanese Patent Application No. 197813/1993 filed on July 15, 1993, Japanese Patent Application No. 253740/1993 filed on September 16, 1993, Japanese Patent Application No. 261875/1993 filed on September 24, 1993 are hereby incorporated by reference.

Other features of the invention will become apparent in the course of the following description of the exemplary embodiments which are given for illustration of the invention and are not intended to be limiting thereof.

EXAMPLES

Various characteristics of the compositions or components thereof prepared in the examples and comparative examples hereinafter are evaluated according to the following methods.

Polystyrene-reduced weight average molecular weight (Mw)

Polystyrene-reduced weight average molecular weights were measured by gel permeation chromatography using a monodispersed polystyrene as a standard, wherein GPC columns (two G2000H $_{\rm XL}$ columns, one G3000H $_{\rm XL}$ column, and one G4000H $_{\rm XL}$ column, all trademarks, manufactured by Tosoh Corp.) were used under the conditions of a flow rate of 1.0 ml/minute, a column temperature of 40°C, and using tetrahydrofuran as an eluant.

Coatability

The chemically amplified resist compositions were applied by spin-coating on 6 inch silicone wafers and baked at 90°C for 2 minutes. Resist coatings thus prepared were observed and those having no unevenness in the coating, being free from cloud and contaminants, and exhibiting high surface smoothness were rated as acceptable.

Photospeed

The exposure dose of irradiation providing 0.5 µm line-and-space patterns as designed (mJ/cm²).

5 Coating reduction

Th yield of the coating reduction was calculated by measuring the resist coating thickness before and after devel pment using an α -step.

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Syntheti Exampi 1

<Synthesis of Resin (A)>

Polyhydroxystyren (30 g) was dissolved in tetrahydrofuran and 10 g of potassium t-butoxise was added to the solution. 60 g of di-t-butyl dicarbonate was added dropwise at 0°C while stirring, and the mixture was reacted for four hours. After the reaction, the resulting solution was dropped into water. The precipitated resin was dried overnight in a vacuum drier at 50°C. The resin thus obtained had a Mw of 15,000. NMR measurement confirmed that the resin has a structure in which 29% of hydrogen atoms in phenolic hydroxy group were replaced by t-butoxycarbonyl group.

Synthetic Example 2

<Synthesis of Resin (A')>

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300 g of p-t-butoxystyrene, 30 g of styrene, and 1.6 g of azobis-iso-butylonitrile were dissolved in 330 g of dioxane, and reacted for 12 hours under a nitrogen atmosphere at 70°C. After the reaction, unreacted monomers were removed to obtain a copolymer of p-hydroxystyrene and styrene. This resin was hydrolyzed with an acid to produce 180 g of a copolymer of p-hydroxystyrene and styrene with Mw of 18,000. NMR measurement confirmed that the molar ratio of p-hydroxystyrene and styrene was 85:15.

Synthetic Example 3

Bisphenol A (15 g) was dissolved in tetrahydrofuran. di-t-butyl dicarbonate in an amount of 2 times and triethylamine in an amount of 0.3 time, respectively the total mols of hydroxy group, were added, and the mixture was reacted for 6 hours under refluxing. After the reaction, the resulting solution was dropped into water. The precipitate was dried overnight in a vacuum drier at 50°C, to obtain dissolution controlling agent (1) of the above-described formula (1).

Synthetic Example 4

A reaction was carried out in the same manner as in Synthetic Example 3 using a compound having hydrogen for the t-butoxycarbonyl group in the dissolution controlling agent of the above-described formula (2) instead of Bisphenol A in Synthetic Example 3, thus obtaining dissolution controlling agent (2) of the formula (2).

Examples 1-17 and Comparative Examples 1-10

Chemically amplified resist compositions were prepared by mixing components shown in Tables 1-1 to 1-3, and removing contaminants by filtering the mixtures through a 0.2 µm filter. These chemically amplified resist compositions were applied to 6 inch silicone wafers by spin coating and baked for 2 minutes at 100°C. Resist coating with a coating thickness of 1 µm was irradiated through a mask using a KrF excimer laser, MBK-400TL-N (trademark, manufactured by Admonscience Co.). After baking for 2 minutes at 110°C, the resist coating was developed for 60 seconds with an aqueous solution of 2.38 wt.% tetramethylammonium hydroxide at 23°C, and rinsed with water, to form resist patterns. Preparation of resist patterns were carried out two times for each chemically amplified resist composition, one immediately after the preparation of the resist composition and the other 30 days after the preparation. The results are shown in Tables 2-1 to 2-3.

Photoacid generators (B), dissolution controlling agents (D), crosslinking agents (E), and solvents (C) in Tables 1-1 to 1-3 are as follows.

Photoacid generators (B)

- (1) triphenylsulfonium trifluoromethanesulfonate
- (2) Methoxyphenyl-bis(trichloromethyl)-s-triazin

Dissolution controlling agent (D)

(1) Compound of formula (1)

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(2) Compound of formula (2)

Crosslinking agent (E)

5 Compound of formula (8)

Solvents (C)

AF: Amyl formate BA: Butyl acetate

ECA: Ethylene glycol monoethylether acetate

EEP: Ethyl-3-ethylpropionate

EL: Ethyl lactate (ethyl 2-hydroxypropionate)

iAA: iso-Amyl acetate

iPB: iso-Propyl butyrate

PGDME: Propylene glycol dimethylether PGEE: Propylene glycol ethylether

PGMEA: Propylene glycol methylether acetate

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Examples Resin Photoacid Controlling Cross Resist 1 (A') (1) - (8) 2 (A') (2) - (8) 3 (A') (2) - (8) Positive-tone Resist 4 (A) (1) 5 (A) (1) 6 (A) (1) 7 (A) (1)	45		35	30	20 ·	10	5
(A') (1) - (A') (1) - (A') (1) (2) - (A') (1) - (A') (2) - (A') (1) (1) - (A') (1) (1) - (A') (1) -				TAB	LE 1-1		
(A') (1) - (A') (2) - (A') (2) - (A') (1) (1) - (A') (1) (1) - (A') (1) (1) - (A') (1) (1) (1) (1) (1) (1) (1) (1) (1) (1	3	Sin	Photoacid generator (B)	Dissolution controlling agent (D)		Crosslinking Solvent (C) agent (E) (Ratio by weight)	Ratio by weight of components
(A') (1) (A') (2) - (A') (2) - (A') (1) (1) - (A') (1) (1) - (A') (1) (1) (1) - (A') (1) (1) (1) (1) (1) (1) (1) (1) (1) (1	ive-tone t						
(A') (2) - (A') (2) - (A) (1) (1) - (A) (1) (1) - (A) (1) (1) (1) (1) (1) (1) (1) (1) (1) (1	<u>ت</u>	((1)	ı	(8)	BA:iPB=50:50	(A):(B):(E):(C) =100:3:35:400
(A') (2) (A) (1) - (A)	3	(, V	(2)	•	(8)	EL:BA:iAA =60:20:20	(A):(B):(E):(C) =100:3:35:400
(A) (1) - (A))	A')	(2)	•	(8)	EL:PGEE:PGMEA =50:20:30	(A):(B):(E):(C) =100:3:35:400
(A) (1) (A) (1) - (A) (1) - (A) (1) (A) (1) (A)	ive-tone t						
(A) (1) - (A) (1) - (A) (1) - (A) (1) - (A)	**	T	(1)	ı		BA=100	(A):(B):(C) =100:3:420
(A) (1)	3	F	(1)		•	EL:BA=50:50	(A):(B):(C) =100:3:420
(1)	S		(1)	1	1	EL:PGMEA=50:50	(A):(B):(C) =100:3:420
	A)		(1)		•	PGMEA=100	(A):(B):(C) =100:3:420

50	45	40	35	30	20	15	5
				TAB	TABLE 1-2		
Examples	Resin (A	A) 9	Photoacid generator (B)	Dissolution controlling agent (D)	Crosslinking agent (E)	Solvent (C) (Ratio by weight)	Ratio by weight of components
∞	(A')		(2)	(1)		BA: AF: 1AA =50:20:30	(A):(B):(D):(C) =100:3:35:400
6	(A')		(1)	(1)	•	EL:iAA=60:40	(A):(B):(D):(C) =100:3:35:400
10	(A')		(2)	(1)		EL:iPB=70:30	(A):(B):(D):(C) =100:3:35:400
, 11	(A')		(2)	(2)	•	EL:iPB=60:40	(A):(B):(D):(C) =100:3:35:400
12	(A')		(2)	(2)	ŧ	EL:BA:AF =50:30:20	(A):(B):(D):(C) =100:3:35:400
13	(A')		(1)	(1)		EL:PGDME=60:40	(A):(B):(D):(C) =100:3:35:400
14	(A')		(1)	(2)	1	EL:PGEE=60:40	(A):(B):(D):(C) =100:3:35:400
15	(A')		(1)	(1)	1	PGMEA:BA=90:10	(A):(B):(D):(C) =100:3:35:400
16	(A')		(1)	(2)	ŧ	PGMEA:EL=90:10	(A):(B):(D):(C) =100:3:35:400
17	(A')		(1)	(2)	J	PGMEA:EEP=90:10	(A):(B):(D):(C) =100:3:35:400

Comparative Examples Resin Negative-tone Resist (A') 2 (A') 3 (A') 3 (A') 4 (A)						
tone (A			TAB	TABLE 1-3		
ative-tone ist		Photoacid generator (B)	Dissolution controlling agent (D)	Crosslinking agent (E)	Solvent (C) (Ratio by weight)	Ratio by weight of components
1 tive-tone						
3 itive-tone ist		(2)		(8)	EL=100	(A):(B):(E):(C) =100:3:35:400
tive-tone		(2)	•	(8)	ECA:BA=50:50	(A):(B):(E):(C) =100:3:35:400
itive-tone ist		(5)	• •	(8)	ECA:EL=50:50	(A):(B):(E):(C) =100:3:35:400
		(1)	1	,	EL=100	(A):(B):(C) =100:3:420
5 (A)		(1)	1	ı	ECA=100	(A):(B):(C) =100:3:420
6 (A')		(1)	(1)	1	EL=100	(A):(B):(D):(C) =100:3:35:400
7 . (A')	-	(1)	(2)	ı	EL=100	(A):(B):(D):(C) =100:3:35:400
8 (A')		(1)	(1)		ECA=100	(A):(B):(D):(C) =100:3:35:400
9 (A')		(1)	(2)	,	ECA:PGDME=50:50	(A):(B):(D):(C) =100:3:35:400
10 (A')		(1)	(2)	•	ECA:PGEE=50:50	(A): (B): (D): (C) 100: 3: 35: 400

	•		1 2							٠		
			30 da		0.04	0.02	0.04		0.03	0.03	0.02	0.02
5		(mm) u	After		· 0	o.	0.		0.	0	0.	0
10	(Coating reduction (µm)	After preparation After 30 days		0.02	0.02	0.04		0.04	0.03	0.02	0.02
15		Coat	After)				Ū	J
20	. 2-1	:m ²).	After preparation After 30 days		20	22	21		59	50	21	20
25	TABLE 2-1	Photospeed (mJ/cm^2) .	tion Af	i								
30		Photospe	ter prepara		20	22	20		28	20	21	20
35		. 4.										
40			Coatability		Acceptable	Acceptable	Acceptable		Acceptable	Acceptable	Acceptable	:
4 5			Examples	ative-tone ist	_	21	8	itive-tone ist	e	S	9	,
50			Exar	Negat Resist		••		Positi Resist	•		_	

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TABLE 2-2

		Photospeed (mJ/cm ²)	J/cm ²)	. Coating reduction (µm)	(wrd) u
Examples	Coatability	After preparation After 30 days	After 30 days	After preparation After 30 days	After 30 days
æ	Acceptable	28	28	0.04	0.04
6	Acceptable	. 25	25	0.04	0.04
10	Acceptable	. 58	. 88	0.03	0.03
11	Acceptable	30	28	0.03	0.03
12	Acceptable	18	18	0.05	90.0
13	Acceptable	31	32	0.05	0.04
14	Acceptable	39	39	0.02	0.05
15	1	32	32	90.0	90.0
16	;	48	48	0.02	0.05
17	:	22	22	0.04	0.03

45 50	40	35	30	25	20	15	10	5		
			-	TABLE 2-3						
Comparative			Photospeed (mJ/cm ²)	mJ/cm²)		Coating	Coating reduction (µm)	(mrt)		I
Examples	Coatability	After	After preparation	After 30	days	After pre	After preparation	After 30	30 days	ys
Negative-tone Resist										1
	Unacceptable		22	14		0.04	4	0.02	25	
2	Unacceptable	•	22	18		0.03	e	0.09	6	
Ю	Acceptable		23	12		0.03	e	0.09	6	
Positive-tone Resist										ı
. 4	Unacceptable		16	10	•	0.03	m	0.03	ლ	
ĸ	Unacceptable		18	17		0.05	ю	0.04	4	
. 9	Unacceptable		34	22		0.05	10	0.23	ю	
7	Unacceptable		46	34		0.02	01	0.08	80	
æ	Unacceptable		37	21		0.13	~	0.0	6	
	Acceptable		38	20		0.05	10	0.16	9	
10	Unacceptable		43	31		0.05		0.14	4	

As fully illustrated above, the compositions for chemically amplified resist of the present invention exhibits superior sensitivity and resolution capability and can be excellently applied especially to large sized substrates by spincoating. The compositions further exhibits excellent storage stability and is suitable as a material for forming chemically amplified resist coating for producing excellently shaped patterns. Because various radiations, such as ultraviolet rays (.g. i-line), deep ultraviolet rays (.g. xcimer laser), X-rays (.g. synchrotron radiation), and charged particle rays (.g. electron beam), are all applicable to the compositions, they can be used with advantage as a chemically amplified resist for the manufacture of semiconductor devices or inte-

grated circuits (ICs) which are expected to be miniaturized in the future.

Obviously, numerous modifications and variations of the pres nt invention are possible in light of the above teachings. It is therefore to be understood that within the scope of the appended claims, the invention may be practiced otherwise than as specifically described herein.

Claims

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- 1. A positive-tone type chemically amplified resist composition comprising,
 - (A) a resin which has an acidic functional group protected by an acid decomposable group and which is converted into an alkali-soluble resin when the acid-decomposable protective group is decomposed,
 - (B) a photoacid generator which is a compound generating an acid by irradiation, and
 - (C) a solvent selected from the group consisting of,
 - (i) an ester of a saturated carboxylic acid having 1-4 carbon atoms with no substituent and an alcohol having 1-6 carbon atoms,
 - (ii) a mixture of ester solvent (i) and an alkyl lactate,
 - (iii) a mixture of an alkyl lactate, a propylene glycol alkyl ether and/or a propylene glycol alkyl ether acetate, and
 - (iv) a propylene glycol alkyl ether acetate.
- 2. The positive-tone resist composition according to claim 1, wherein the amount of said acid-decomposable protective groups introduced to said component (A) is 15-100% of the total number of the acidic functional groups and the acid-decomposable protective groups.
- The positive-tone resist composition according to claim 1, wherein 0.05-20 parts by weight of component (B) and 20-3,000 parts by weight of component (C) are incorporated for 100 parts by weight of component (A).
 - 4. A positive-tone type chemically amplified resist composition comprising,
 - (A') an alkali-soluble resin,
 - (D) a compound which inhibits dissolution of the alkali-soluble resin in alkali and is decomposable in the presence of an acid, and, when decomposed in the presence of an acid, exhibits an action of reducing or losing the alkali insolubility of a resin coating, or accelerating dissolution of the alkali-soluble resin in alkali
 - (B) a photoacid generator which is a compound generating an acid by irradiation, and
 - (C) a solvent selected from the group consisting of,
 - (i) an ester of a saturated carboxylic acid having 1-4 carbon atoms with no substituent and an alcohol having 1-6 carbon atoms,
 - (ii) a mixture of ester solvent (i) and an alkyl lactate,
 - (iii) a mixture of an alkyl lactate, a propylene glycol alkyl ether and/or a propylene glycol alkyl ether acetate, and
 - (iv) a propylene glycol alkyl ether acetate.
 - The positive-tone resist composition according to claim 4, wherein 0.05-20 parts by weight of component (B), 20-3,000 parts by weight of component (C), 5-150 parts by weight of component (D) are incorporated for 100 parts by weight of component (A').
 - 6. A negative-tone type chemically amplified resist composition comprising,
 - (A') an alkali-soluble resin,
 - (E) a compound which can crosslink the alkalisoluble resin in the presence of an acid,
 - (B) a photoacid generator, and
 - (C) a solvent selected from the group consisting of,
 - (i) an ester of a saturated carboxylic acid having 1-4 carbon atoms with no substituent and an alcohol having 1-6 carbon atoms,
 - (ii) a mixture of ester solvent (i) and an alkyl lactate, and
 - (iii) a mixture of an alkyl lactate, a propyl n glycol alkyl ether and/or a propyl ne glycol alkyl ether acet-
 - 7. The negative-ton resist composition according to claim 7, wh rein 5-95 parts by w ight of compon nt

(E), 0.05-20 parts by weight of component (B), and 20-3,000 parts by w ight of component (C) are incorporated for 100 parts by weight of component (A').



EUROPEAN SEARCH REPORT

Application Number P 94 30 5148

Category	Citation of document with in of relevant pa	ndication, where appropriate,	Relevant to chain	CLASSIFICATION (In APPLICATION (In	OF THE
X	EP-A-0 388 343 (INT MACHINES)	ERNATIONAL BUSINESS	1-5	G03F7/004	
	* page 4, line 50 -	line 53; examples			
Y	5,7-9 * * the whole documen	t *	1-7		
Y	EP-A-0 425 411 (INT MACHINES)	ERNATIONAL BUSINESS	1-7		
		line 31; claim 12 *			
Y	EP-A-0 440 374 (WAK	D PURE CHEMICALS IND.	1-7		
	* page 8, line 58 -	page 9, line 6 *			
Y	EP-A-0 510 670 (SUM * page 6 - page 7 *	ITOMO CHEMICAL CO LTD)	1-7	·	
Y	EP-A-0 540 965 (BAS * page 7, line 6 -	 F AG) line 15 *	1-7		
Y	WO-A-92 08166 (INTERMACHINES)	RNATIONAL BUSINESS	1-7	TECHNICAL PIE SEARCHED (LDS bitCL6)
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X : part Y : part doc	CATEGORY OF CITED DOCLIMEN featurity relevant if taken alone icolarly relevant if combined with and most of the same category	E : earlier paient do after the filles d	ato In the application		
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